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AMENDMENTS

Amendments to the Claims:

The currently pending and amended claims are below. Please amend the claims following, wherein the deleted matter is shown by strikethrough and the added matter is shown by underlining.

- (Previously presented) A nanoparticle probe composition for facilitating molecular imaging or monitoring, comprising:
 - a detectable moiety comprising a magnetic nanoparticle having a biocompatible coating thereon:
 - a targeting probe attached to the biocompatible coating, wherein the targeting probe is selected from the group consisting of a nucleic acid probe, an antibody, an antibody fragment, and an aptamer; and
 - an intracellular delivery ligand attached to the biocompatible coating.
- 2. (Withdrawn) The composition of Claim 1, wherein the targeting probe is a nucleic acid probe.
- 3. (Withdrawn) The composition of Claim 2, wherein the nucleic acid probe hybridizes to a nucleic acid target sequence on a subject nucleic acid and forms a stem-loop structure when not bound to the nucleic acid target sequence.
- 4. (Withdrawn) The composition of Claim 2, wherein the nucleic acid probe comprises a modification of the nucleic acid backbone for the increased stability of the nucleic acid as compared to a naturally occurring nucleic acid.
- (Canceled)
- (Previously presented) The composition of Claim 1, wherein the targeting probe is an antibody or fragment thereof.
- 7. (Withdrawn) The composition of Claim 1, wherein the targeting probe is an aptamer.

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8. (Original) The composition of Claim 1, wherein the composition comprises two or more

different targeting probes, wherein each of the two or more targeting probes is selected from the group consisting of a nucleic acid probe, a polypeptide probe, an antibody, an antibody fragment,

group consisting of a nacione acid proce, a polypeptace proce, an analous, an analous, magnet

a high affinity ligand, a peptide, and an aptamer.

9. (Previously presented) The composition of Claim 1, wherein the intracellular delivery ligand

comprises a protein transduction peptide selected from the group consisting of HIV-1 TAT, HSV

VP22, and ANTP.

10. (Withdrawn) The composition of Claim 1, wherein the intracellular delivery ligand is selected from the group consisting of the W/R peptide (SEQ ID NO:4), the NLS* peptide (SEQ

ID NO:5), the AlkCWK18 peptide (SEQ ID NO:6), DiCWK18 peptide (SEQ ID NO:7), the

transportan peptide (SEQ ID NO:8), the DipaLytic peptide (SEQ ID NO:9), the K16RGD peptide

(SEQ ID NO:10), the P1 peptide (SEQ ID NO:11), the P2 peptide (SEQ ID NO:12), the P3

peptide (SEQ ID NO:13), the P3a peptide (SEQ ID NO:14), the P9.3 peptide (SEQ ID NO:15), the Plae peptide (SEO ID NO:16), the Kplae peptide (SEQ ID NO:17), the cKplae peptide (SEQ

ID NO:18), the MGP peptide (SEQ ID NO:19), the HA2 peptide (SEQ ID NO:20), the LARL46

peptide (SEQ ID NO:21), the Hel-11-7 peptide (SEQ ID NO:22), the KK peptide (SEQ ID

NO:23), the KWK peptide (SEQ ID NO:24), the RWR peptide (SEQ ID NO:25), and the

Loligomer peptide (SEQ ID NO:26).

11. (Previously presented) The composition of Claim 1, wherein the intracellular delivery ligand

facilitates receptor-mediated endocytosis of the composition.

12. (Previously presented) The composition of Claim 1, wherein the intracellular delivery ligand

facilitates entry into a cell by permeabilizing the cell membrane.

13. (Original) The composition of Claim 1, wherein the magnetic nanoparticle comprises a metal

detectable by use of an MRI instrument that is selected from the group consisting of selected from the group consisting of iron, cobalt, zinc, cadmium, nickel, gadolinium, chromium, copper,

manganese, terbium, europium, gold, silver, platinum, and alloys thereof,

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14. (Original) The composition of Claim 13, wherein the magnetic nanoparticle is selected from

the group consisting of monocrystalline iron oxide nanoparticle (MION), chelate of gadolinium,

and superparamagnetic iron oxide (SPIO).

15. (Original) The composition of Claim 13, wherein the magnetic nanoparticle is

monocrystalline iron oxide nanoparticle (MION).

16. (Original) The composition of Claim 13, wherein the magnetic nanoparticle is a free metal

ion, a metal oxide, a chelate, or an insoluble metal compound.

17. (Original) The composition of Claim 13, wherein the magnetic nanoparticle is selected from

the group consisting of Fe₃O₄, Fe₂O₄, Fe_xPt_y, Co_xPt_y, MnFe_xO_y, CoFe_xO_y, NiFe_xO_y, CuFe_xO_y,

ZnFe_xO_y, and CdFe_xO_y, wherein x and y vary between 1 and 6, depending on the method of

synthesis.

18. (Original) The composition of Claim 13, wherein the magnetic nanoparticle further comprises a metal coating selected from the group consisting of gold, silver, iron, cobalt, zinc.

cadmium, nickel, gadolinium, chromium, copper, and manganese, and an alloy thereof.

19. (Original) The composition of Claim 1, wherein the biocompatible coating is selected from

the group consisting of a surfactant based coating, a starch based coating, a dextran based

coating, a silica based coating, a layer by layer coating, a phospholipid-polyethylene glycol

coating, a polymer coating, a mesoporous particle coating, a microporous particle coating, a lipid

based coating, and a dendrimer based coating.

20. (Original) The composition of Claim 1, wherein the biocompatible coating is a

phospholipid-polyethylene glycol coating.

21. (Original) The composition of Claim 1, wherein the composition further comprises a second

delivery ligand that is attached to the biocompatible coating and that interacts with a molecule

located on the outer surface of a particular type of cell or tissue.

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22. (Original) The composition of Claim 21, wherein the second delivery ligand is selected from

a group consisting of an antibody, an antibody fragment, a peptide, an aptamer, a receptor-

specific ligand, and a tissue-specific ligand.

23. (Original) The composition of Claim 21, wherein the second delivery ligand that interacts

with an infected cell or a diseased cell.

24. (Previously presented) The composition of Claim 1, further comprising a second detectable

moiety attached to the biocompatible coating, wherein the second detectable moiety is selected

from the group consisting of a resonance energy transfer donor mojety; a resonance energy

transfer acceptor moiety; a radioisotope; a fluorescent dye; an organic bead, a chelator, or a

magnetic nanoparticle with fluorescent or luminescent characteristics; and an inorganic bead with

fluorescent or luminescent characteristics.

25. (Original) The composition of Claim 1, further comprising a therapeutic molecule attached

to the biocompatible coating or to the magnetic nanoparticle.

26. (Previously presented) A composition for facilitating signal transduction in molecular

imaging or monitoring, comprising:

a first magnetic nanoparticle probe composition comprising a detectable moiety

comprising a magnetic nanoparticle having a biocompatible coating thereon; a first targeting probe attached to the biocompatible coating; and an intracellular delivery ligand

attached to the biocompatible coating; and

a second magnetic nanoparticle probe composition comprising a detectable moiety

comprising a magnetic nanoparticle having a biocompatible coating thereon; a second

targeting probe attached to the biocompatible coating; and an intracellular delivery ligand

attached to the biocompatible coating;

wherein the first targeting probe binds to a first target and the second targeting probe binds with a second target; and wherein an effect on water relaxation from interaction between the first and

second magnetic nanoparticles can be detected to determine binding of both the first and the

second targeting probes to the first and the second target, and wherein the first magnetic

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nanoparticle probe composition is different than the second magnetic nanoparticle probe

composition.

27. (Withdrawn) The composition of Claim 26, wherein the first targeting probe and the second

targeting probe are nucleic acid probes; wherein the first nucleic acid probe hybridizes to a first

nucleic acid target sequence on a subject nucleic acid and the second nucleic acid probe

hybridizes with a second nucleic acid target sequence on the subject nucleic acid; and wherein

the first nucleic acid target sequence and the second nucleic acid target sequence are separated by a number of nucleotides on the subject nucleic acid such that an effect on water relaxation from

interaction between the first and second magnetic nanoparticles can be detected to determine

hybridization of both the first and the second nucleic acid probes.

28. (Withdrawn) The composition of Claim 27, further comprising

a third magnetic nanoparticle probe composition comprising a detectable moiety

comprising a magnetic nanoparticle having a biocompatible coating thereon; a third

targeting probe attached to the biocompatible coating; and an intracellular delivery ligand attached to the biocompatible coating; and

a fourth magnetic nanoparticle probe composition comprising a detectable moiety

comprising a magnetic nanoparticle having a biocompatible coating thereon; a fourth

targeting probe attached to the biocompatible coating; and an intracellular delivery ligand

attached to the biocompatible coating;

wherein the third nucleic acid probe hybridizes to a third nucleic acid target sequence on the subject nucleic acid and the fourth nucleic acid probe hybridizes with a fourth nucleic acid target

sequence on the subject nucleic acid; and wherein the third nucleic acid target sequence and the

fourth nucleic acid target sequence are separated by a number of nucleotides on the subject

nucleic acid such that an effect on water relaxation from interaction between the third and fourth

magnetic nanoparticles can be detected to determine hybridization of both the third and the

fourth nucleic acid probes.

29. (Withdrawn) The composition of Claim 27, wherein the composition further comprises

additional magnetic nanoparticle probe pairs.

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30. (Withdrawn) The composition of Claim 27, wherein the first nucleic acid probe, the second

nucleic acid probe, or both nucleic acid probes hybridize to a nucleic acid target sequence on a

subject nucleic acid and form a stem-loop structure when not bound to the nucleic acid target

sequence.

31. (Withdrawn) The composition of Claim 27, wherein the first nucleic acid probe, the second

nucleic acid probe, or both nucleic acid probes comprise a modification of the nucleic acid backbone for the increased stability of the nucleic acid as compared to a naturally occurring

nucleic acid.

32. (Original) The composition of Claim 26, wherein the first and the second targeting probes

are polypeptide probes; wherein the first polypeptide probe binds to a first target sequence on a subject polypeptide and the second polypeptide probe binds with a second target sequence on a

subject polypeptide; and wherein the first target sequence and the second target sequence are

separated by a distance such that an effect on water relaxation from interaction between the first

and second magnetic nanoparticles can be detected to determine binding of both the first and the

second polypeptide probes.

33. (Original) The composition of Claim 32, wherein the first and the second target sequences

are located on a single subject polypeptide.

34. (Original) The composition of Claim 32, wherein the first target sequence is located on a

first subject polypeptide and the second target sequence is located on a second subject polypeptide; and wherein effect on water relaxation from interaction between the first and second

magnetic nanoparticles can be detected to determine the interaction of the first subject

magnetic nanoparticles can be detected to determine the interaction of the first subject

polypeptide and the second subject polypeptide.

35. (Original) The composition of Claim 32, wherein the first targeting probe, the second

targeting probe, or both targeting probes are antibodies or fragments thereof.

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36. (Withdrawn) The composition of Claim 26, wherein the first targeting probe, the second targeting probe, or both targeting probes are selected from the group consisting of a high affinity ligand, a peptide, and an aptamer.

37. (Original) The composition of Claim 26, wherein the first or the second magnetic nanoparticle probe composition or both comprise two or more targeting probes.

38. (Previously presented) The composition of Claim 26, wherein the intracellular delivery ligand comprises a cell penetrating peptide selected from the group consisting of HIV-1 TAT, HSV VP22, and ANTP.

39. (Withdrawn) The composition of Claim 26, wherein the intracellular delivery ligand is selected from the group consisting of the W/R peptide (SEQ ID NO:4), the NLS* peptide (SEQ ID NO:5), the AlkCWK18 peptide (SEQ ID NO:6), the DiCWK18 peptide (SEQ ID NO:7), the transportan peptide (SEQ ID NO:8), the DipaLytic peptide (SEQ ID NO:9), the K16RGD peptide (SEQ ID NO:10), the P1 peptide (SEQ ID NO:11), the P2 peptide (SEQ ID NO:12), the P3 peptide (SEQ ID NO:13), the P3a peptide (SEQ ID NO:14), the P9.3 peptide (SEQ ID NO:15), the Plae peptide (SEQ ID NO:16), the Kplae peptide (SEQ ID NO:17), the cKplae peptide (SEQ ID NO:18), the MGP peptide (SEQ ID NO:19), the HA2 peptide (SEQ ID NO:20), the LARL46 peptide (SEQ ID NO:21), the Hel-11-7 peptide (SEQ ID NO:22), the KK peptide (SEQ ID NO:23), the KWK peptide (SEQ ID NO:24), the RWR peptide (SEQ ID NO:25), and the Loligomer peptide (SEQ ID NO:26).

40. (Original) The composition of Claim 26, wherein the magnetic nanoparticle comprises a metal detectable by use of an MRI instrument that is selected from the group consisting of selected from the group consisting of iron, cobalt, zinc, cadmium, nickel, gadolinium, chromium, copper, manganese, terbium, europium, gold, silver, platinum, and alloys thereof.

41. (Original) The composition of Claim 40, wherein the magnetic nanoparticle is monocrystalline iron oxide nanoparticle (MION).

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42. (Original) The composition of Claim 40, wherein the magnetic nanoparticle is a free metal

ion, a metal oxide, a chelate, or an insoluble metal compound.

43. (Previously presented) The composition of Claim 40, wherein the magnetic nanoparticle is

selected from the group consisting of Fe₃O₄, Fe₂O₄, Fe_xPt_y, Co_xPt_y, MnFe_xO_y, CoFe_xO_y, NiFe_xO_y,

 $\label{eq:cuFexOy} \text{CuFe}_x O_y, \ \text{ZnFe}_x O_y, \ \text{and} \ \text{CdFe}_x O_y, \ \text{wherein} \ x \ \text{and} \ y \ \text{vary} \ \text{between} \ 1 \ \text{and} \ 6 \ \text{depending} \ \text{on} \ \text{the}$

method of synthesis.

44. (Original) The composition of Claim 40, wherein the magnetic nanoparticle further

comprises a metal coating selected from the group consisting of gold, silver, iron, cobalt, zinc,

cadmium, nickel, gadolinium, chromium, copper, manganese, and an alloy thereof.

45. (Original) The composition of Claim 26, wherein the biocompatible coating is selected from

the group consisting of a surfactant based coating, a starch based coating, a dextran based

coating, a silica based coating, a layer by layer coating, a phospholipid-polyethylene glycol coating, a polymer coating, a microporous particle coating, a lipid

based coating, and a dendrimer based coating.

46. (Original) The composition of Claim 26, wherein the biocompatible coating is a

phospholipid-polyethylene glycol coating.

47. (Original) The composition of Claim 26, wherein the first magnetic nanoparticle probe

composition, the second magnetic nanoparticle probe composition, or both further comprise a second delivery ligand that is attached to the biocompatible coating and that interacts with a

molecule located on the outer surface of a particular type of cell or tissue.

48. (Original) The composition of Claim 47, wherein the second delivery ligand is selected from

a group consisting of an antibody, an antibody fragment, a peptide, an aptamer, a receptor-

specific ligand, and a tissue-specific ligand.

49. (Previously presented) The composition of Claim 47, wherein the second delivery ligand

interacts with an infected cell or a diseased cell.

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50. (Original) The composition of Claim 26, wherein the first magnetic nanoparticle probe

composition, the second magnetic nanoparticle probe composition, or both further comprise a second detectable moiety attached to the biocompatible coating, wherein the detectable moiety is

selected from the group consisting of a resonance energy transfer donor or acceptor moiety, a

fluorescent dye, an organic bead with fluorescent or luminescent characteristics, and an inorganic

bead with fluorescent or luminescent characteristics.

51. (Original) The composition of Claim 26, wherein the first magnetic nanoparticle probe

composition, the second magnetic nanoparticle probe composition, or both further comprise a

therapeutic molecule attached to the biocompatible coating or to the magnetic nanoparticle.

52. (Original) A composition for facilitating molecular imaging, comprising two or more

magnetic nanoparticle probe compositions within a vesicle, wherein each magnetic nanoparticle

probe composition comprises at least one targeting probe and a detectable moiety attached

thereto, wherein the detectable moiety comprises a magnetic nanoparticle having a biocompatible coating thereon, and wherein the vesicle comprises a biocompatible membrane having at least

coating thereon, and wherein the vesicle comprises a biocompatible membrane

one delivery ligand on its outer surface.

Claims 53.-86.(Canceled)

87. (Previously presented) A method for producing a magnetic nanoparticle probe composition

for facilitating molecular imaging, comprising:

combining a magnetic nanoparticle with a biocompatible coating;

adding an intracellular delivery ligand; and

adding a targeting probe, wherein the targeting probe is selected from the group

consisting of a nucleic acid probe, an antibody, an antibody fragment, and an aptamer.

88. (Previously presented) The method of Claim 87, wherein the intracellular delivery ligand is selected from the group consisting of the HIV-1 TAT peptide (SEO ID NO:1), the HSV VP22

peptide (SEQ ID NO:2), the ANTP peptide (SEQ ID NO:3), the W/R peptide (SEQ ID NO:4),

the NLS* peptide (SEQ ID NO:5), the AlkCWK18 peptide (SEQ ID NO:6), the DiCWK18

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peptide (SEQ ID NO:7), the transportan peptide (SEQ ID NO:8), the DipaLytic peptide (SEQ ID

NO:9), the K16RGD peptide (SEQ ID NO:10), the P1 peptide (SEQ ID NO:11), the P2 peptide

(SEQ ID NO:12), the P3 peptide (SEQ ID NO:13), the P3a peptide (SEQ ID NO:14), the P9.3 $\,$

peptide (SEQ ID NO:15), the Plae peptide (SEQ ID NO:16), the Kplae peptide (SEQ ID NO:17),

the cKplae peptide (SEQ ID NO:18), the MGP peptide (SEQ ID NO:19), the HA2 peptide (SEQ ID NO:20), the LARL46 peptide (SEO ID NO:21), the Hel-11-7 peptide (SEO ID NO:22), the

KK peptide (SEQ ID NO:23), the KWK peptide (SEQ ID NO:24), the RWR peptide (SEQ ID

NO:25), and the Loligomer peptide (SEQ ID NO:26).

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89. (Original) The method of Claim 87, wherein the magnetic nanoparticle comprises a metal

detectable by an MRI instrument that is selected from the group consisting of selected from the group consisting of iron, cobalt, zinc, cadmium, nickel, gadolinium, chromium, copper, and

manganese.

90. (Original) The method of Claim 87, wherein the magnetic nanoparticle is monocrystalline

iron oxide nanoparticle (MION).

91. (Original) The method of Claim 87, wherein the biocompatible coating is selected from the

group consisting of a surfactant based coating, a starch based coating, a dextran based coating, a

silica based coating, a layer by layer coating, a phospholipid-polyethylene glycol coating, a polymer coating, a mesoporous particle coating, a microporous particle coating, a lipid based

coating, and a dendrimer based coating.

92. (Original) The method of Claim 87, wherein the biocompatible coating is a phospholipid-

polyethylene glycol coating.

93. (Previously presented) The method of Claim 87, further comprising the step of adding a

second delivery ligand to the biocompatible coating that interacts with a molecule located on the

outer surface of a particular type of cell or tissue, wherein the second delivery ligand is selected

from a group consisting of an antibody, an antibody fragment, a peptide, an aptamer, a receptor-

specific ligand, and a tissue-specific ligand.

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94. (Original) The method of Claim 87, further comprising the step of adding a therapeutic molecule to the biocompatible coating.

95. (Withdrawn) A nanoparticle probe composition for facilitating molecular imaging or monitoring, comprising:

a detectable moiety comprising a magnetic nanoparticle having a biocompatible coating thereon:

a targeting probe attached to the biocompatible coating; and

an intracellular delivery ligand attached to the biocompatible coating,

wherein the intracellular delivery ligand is selected from the group consisting of the HSV VP22 peptide (SEQ ID NO:2), the ANTP peptide (SEQ ID NO:3), the W/R peptide (SEQ ID NO:4), the NLS* peptide (SEQ ID NO:5), the AlkCWK18 peptide (SEQ ID NO:6), the DiCWK18 peptide (SEQ ID NO:7), the transportan peptide (SEQ ID NO:8), the DipaLytic peptide (SEQ ID NO:9), the K16RGD peptide (SEQ ID NO:10), the P1 peptide (SEQ ID NO:11), the P2 peptide (SEQ ID NO:12), the P3 peptide (SEQ ID NO:13), the P3a peptide (SEQ ID NO:14), the P9.3 peptide (SEQ ID NO:15), the P1ae peptide (SEQ ID NO:16), the Kplae peptide (SEQ ID NO:17), the cKplae peptide (SEQ ID NO:18), the MGP peptide (SEQ ID NO:19), the HA2 peptide (SEQ ID NO:20), the LARL46 peptide (SEQ ID NO:21), the Hel-11-7 peptide (SEQ ID NO:22), the KK peptide (SEQ ID NO:23), the KWK peptide (SEQ ID NO:24), the RWR peptide (SEQ ID NO:25), and the Loligomer peptide (SEQ ID NO:26).

96. (Withdrawn) A method for producing a magnetic nanoparticle probe composition for facilitating molecular imaging, comprising:

combining a magnetic nanoparticle with a biocompatible coating;

adding a targeting probe; and

adding an intracellular delivery ligand;

wherein the intracellular delivery ligand is selected from the group consisting of the HSV VP22 peptide (SEQ ID NO:2), the ANTP peptide (SEQ ID NO:3), the W/R peptide (SEQ ID NO:4), the NLS* peptide (SEQ ID NO:5), the AlkCWK18 peptide (SEQ ID NO:6), the DicWK18 peptide (SEQ ID NO:7), the transportan peptide (SEQ ID NO:8), the DipaLytic peptide (SEQ ID NO:8).

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NO:9), the K16RGD peptide (SEQ ID NO:10), the P1 peptide (SEQ ID NO:11), the P2 peptide (SEQ ID NO:12), the P3 peptide (SEQ ID NO:13), the P3a peptide (SEQ ID NO:14), the P9.3 peptide (SEQ ID NO:15), the Plae peptide (SEQ ID NO:16), the Kplae peptide (SEQ ID NO:17), the cKplae peptide (SEQ ID NO:18), the MGP peptide (SEQ ID NO:19), the HA2 peptide (SEQ ID NO:20), the LARL46 peptide (SEQ ID NO:21), the Hel-11-7 peptide (SEQ ID NO:22), the KK peptide (SEQ ID NO:23), the KWK peptide (SEQ ID NO:24), the RWR peptide (SEQ ID NO:25), and the Loligomer peptide (SEQ ID NO:26).